DEC 1 3 2004

510(k) Summary for TERATECH Model 8IOC4, 8IOL4, and 10LAP4 Probes

1. Sponsor

Terratech Corporation 77-79 Terrace Hall Rd. Burlington, MA 01803

Contact Person:

Charles F. Hottinger, Ph.D., RAC,

Regulatory Affairs Consultant

Telephone:

206-780-7945

Date Prepared:

November 5, 2004

2. DEVICE NAME

Proprietary Name:

TERATECH Model 8IOC4, 8IOL4, and 10LAP4

Probes

Common/Usual Name:

Diagnostic Ultrasound Transducer

Classification Name:

Diagnostic Ultrasound Transducer

(21 CFR 892.1570, 90-ITX)

3. PREDICATE DEVICES

Subject Device	Predicate 1	Predicate 2
8IOL4	Philips LI9-5	TERATECH 10V5
8IOC4	Philips CT8-4	TERATECH 10V5
10LAP4	Philips LAP L9-5	

The Philips probes are marketed for use with the Philips HDI 5000; that system has been cleared in following 510(k) submissions, among possibly others: K961459, K991671, K994373, K002003, and K011224.

The TERATCH predicate probes, as well as the subject devices, are used with the TERATECH Model 2000 portable imaging system. This system has been cleared under the following 510(k) submissions: K992595,

K010883, K012191, K030191, and K040840.

4. INTENDED USE

The TERATECH Model 8IOC4, 8IOL4, and 10LAP4 Probes are intended for diagnostic ultrasound imaging or fluid flow analysis of the human body; specific indications for use a tabulated in Section 4.3 of this submission.

5. DEVICE DESCRIPTION

The TERATECH Model 8IOC4, 8IOL4, and 10LAP4 Probes are intended for use with the Model TERATECH2000, a portable ultrasoundimaging system. Technical specifications for the Model 8IOC4, 8IOL4, and 10LAP4 Probes with the Model 2000 are as follows:

Model	8IOC4	8IOL4	10LAP4
Frequency/	6.0 MHz	7.5 MHz	7.0 MHz
# Elements	128	128	128
Апау type	Curved	Linear	Linear
Pitch (mm)	0.32	0.30	0.30
Elevation width (mm		5.0	5.0
Geometric focus (mm)		25	25
Azimuth radius (mm	40	N/A	N/A
Azimuth length (mm	50.0	38.4	38.4

6. BASIS FOR SUBSTANTIAL EQUIVALENCE

The TERATECH Model 8IOC4, 8IOL4, and 10LAP4 Probes are substantially equivalent to the above cited Philips transducers, which are currently in commercial distribution in the United States. The TERATECH Model 8IOC4, 8IOL4, and 10LAP4 Probes are believed to be identical in mechanical design and materials to the respective Philips, and are intended for the same clinical applications.

4.3 INDICATIONS FOR USE

The TERATECH Model 8IOC4, 8IOL4, and 10LAP4 Probes are intended for the uses described in the Diagnostic Ultrasound Indications For Use Form is provided below.



Food and Drug Administration 9200 Corporate Boulevard Rockville MD 20850

DEC 1 3 2004

TERATECH Corporation % Mr. Mark Job Responsible Third Party Official Regulatory Technology Services 1394 25th Street NW BUFFALO MN 55313

Re: K043278

Trade Name: Terason (Teratech) Model 2000 Portable Ultrasound System and

Teratech Model 8IOC4, 8IOL4, 10LAP4 Probes

Regulation Number: 21 CFR 892.1550

Regulation Name: Ultrasonic pulsed Doppler imaging system

Regulation Number: 21 CFR 892.1560

Regulation Name: Ultrasonic pulsed echo imaging system

Regulation Number: 21 CFR 892.1570

Regulation Name: Diagnostic ultrasonic transducer

Regulatory Class: II

Product Code: 90 IYN, IYO, and ITX

Dated: November 22, 2004 Received: November 26, 2004

Dear Mr. Job:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and we have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

This determination of substantial equivalence applies to the following transducers intended for use with the Terason (Teratech) Model 2000 Portable Ultrasound System, as described in your premarket notification:

Transducer Model Number

Teratech Model 8IOC4
Teratech Model 8IOL4
Teratech Model 10LAP4

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act);-21 CFR 1000-1050.

This determination of substantial equivalence is granted on the condition that prior to shipping the first device, you submit a postclearance special report. This report should contain complete information, including acoustic output measurements based on production line devices, requested in Appendix G, (enclosed) of the Center's September 30, 1997 "Information for Manufacturers Seeking Marketing Clearance of Diagnostic Ultrasound Systems and Transducers." If the special report is incomplete or contains unacceptable values (e.g., acoustic output greater than approved levels), then the 510(k) clearance may not apply to the production units which as a result may be considered adulterated or misbranded.

The special report should reference the manufacturer's 510(k) number. It should be clearly and prominently marked "ADD-TO-FILE" and should be submitted in duplicate to:

Food and Drug Administration Center for Devices and Radiological Health Document Mail Center (HFZ-401) 9200 Corporate Boulevard Rockville, Maryland 20850

This letter will allow you to begin marketing your device as described in your premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus permits your device to proceed to market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Office of Compliance at (240) 276-0120. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 443-6597 or at its Internet address http://www.fda.gov/cdrh/dsma/dsmamain.html

If you have any questions regarding the content of this letter, please contact Rodrigo C. Perez at (301) 594-1212.

Sincerely yours,

Nancy C. Brogdon

Director, Division of Reproductive, Abdominal and Radiological Devices

Office of Device Evaluation

Center for Devices and Radiological Health

Enclosure(s)

Terason Model 2000 Portable Ultrasound System System:

Transducer: (see comments) Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Diagnostic ultrasound imaging	DI NUIC	of Ope	colion		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
Clinical Applica General (Track I Only)	Specific (Tracks I & III)	6	M	PWD	CWD	Calor Dopp*	Comb. Modes	Other
Ophthalmic	Ophthalmic							
Оргинание	Fetai	P ^{1/4}	PX	P".	Γ	P ^{z,i}	PZA	Pz
	Abdominal*:	P''	P ^{Z4}	PZA		Pzi	Pzi	PZ
	Intra-operative (Spec.)	P	P*	P'	T	P.	P'	P
	Intra-operative (Neuro)	P ³	P ₂	P'	1	P3	P'	P.
	Laparoscopic	N	IN	N	I	N	N	N.
	Pediatric :	PIA	P44	P ^{2/2}		PIA	PZA	P**
Fetal Imaging	Small Organ (Thyroid,	PZA	P	Pir		P ¹³	Pz	PZI
& Other	Breast, Testes, etc.):	Pur	PZA	PW	1	PZA	PZI	Pza
	Neonatal Cephalica:	FIF	PIA	PZ		PZA	P ^{2,4}	P
	Adult Cephalic*:	PA	PP	부 px -		P SM	PH	PH
	Trans-rectal:	P	P ³⁴	PH	 	P	PH	P
	Trans-vaginal.	╂╌╌╌		 	 		T	
	Trans-urethral		+	1	 	1		T
	Trans-esoph. (non-Card.)	PZT	P2.1	427	 	PZ	P ^{z,1}	P2.4
	Musculo-skel, (Convent.):	Pitt	PZE	PZZ	1 —	P	P4	P'1
	Musculo-skel. (Superfic)	 '	+'	 	1	1		
	Intra-luminal	1		1	1	T -		
	Other (Specify)	P	P ²	P ²	1	P'	P ²	P
Cardiac	Cardiac Adult	P -	P	\ -	 	P'	P'	P ²
	Cardiac Pediatric	 - -	- -	+	 	 	 	1
	Trans-esoph. (Cardiac)	 -	 	+-		 		1
	Other (Specify)			P2.4	 	P2.7	PZ	PZA
Peripheral	Peripheral vessel ^a :	P ₁.t	P ²⁴	P		1 P	+	+
Vessel	Other (Specify)	<u> </u>			A			┸——

N= new indication; P= previously cleared by FDA; E= added under Appendix E
Includes Color Doppler (CD), Directional Power Doppler (DPD), and (non-directional) Power Doppler.

*8+M; B+PWD; B+CD; B+DPD; B+PD.

*Harmonic Imaging (HI)
Includes ultrasound guidance for placement of needles, cetheters.
Abdominal organs and peripheral vessel.

Abdominal organs and peripheral vessel.

Includes ultrasound guidance for placement of needles, catheters, cryosurgery, and brachytherapy Includes ultrasound guidance of transvaginal biopsy, infertifity monitoring of folicle development.

Includes guidance of amniocentesis, infertifity monitoring of folicle development.

System uses previously cleared under K992505 with 3 MHz Model L3 (Linear).

System uses previously cleared under K012191.

System uses previously cleared under K010883.

System uses previously cleared under K030191.

System uses previously cleared under K0301840.

System uses previously cleared under KO40840.

System uses previously cleared under KO40840.
Includes uses in military field settings in addition to hospital/clinic settings.
(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of Center for Devices and Radiological Health, Office of Device Evaluation Prescription Use (Per 21 CFR 801,109)

(Division Sign-Off)

Division of Reproductive, Abdominal

and Radiological Devices

510(k) Number _

TERATECH Corp. 510(k)

12/09/04

Teratech Model 8IOC4, 81OL4, and 10LAP4 Prohes

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System:	Tereson Mode	2000 Portable Ultrasqun	d System
Transducer:	810C4		

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows: Mode of Operation Clinical Application CWD Other Color Comb. Specific В М PWD General Dopp* Modes (Track I Only) (Tracks I & III) Ophthalmic Ophthalmic Fetal* Abdominal* N N, N, N Intra-operative (Spec.) Intra-operative (Neuro) Laperoscopic Pediatric : Fetal Small Organ (Thyroid, Imaging Breast, Testes, etc.)' Neonatal Cephalic*: & Other Adult Cephalic : Trans-rectal": Trans-vaginal3: Trans-urethral Trans-esoph. (non-Card.) Musculo-skel. (Convent.) Musculo-skel, (Superfic) Intra-luminal Other (Specify) Cardiac Adult Cardiac Cardiac Pediatric Trans-esoph. (Cardiac) Other (Specify) Peripheral vessel Peripheral Other (Specify) Vessel

N= new indication; P= previously cleared by FDA; E= added under Appendix E

Includes Color Doppler (CD), Directional Power Doppler (DPO), and (non-directional) Power Doppler

B+M; B+PWD; B+CD; B+DPD; B+PD.

Harmonic Imaging (HI)

Includes ultrasound guidance for placement of needles, catheters.

Abdominal organs and peripheral vessel.

Includes ultrasound guidance for placement of needles, catheters, cryosurgery, and brachytherapy Includes ultrasound guidance of transvaginal blopsy, infentility monitoring of follicle development.

*Includes ultrasound guidance of variovaginal olopsy, menting monitoring of folicle development.

*Includes guidance of amniocentesis, infertility monitoring of folicle development.

Additional Comments: P1: uses previously cleared under K992505 with 3 MHz Model L3 (Linear);

P1: uses previously cleared under K012191; P3: uses previously cleared under K010883; P4: uses previously cleared

Includes uses in military field settings in addition to hospital/clinic settings.
(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of Center for Devices and Radiological Health, Office of Device Evaluation Prescription Use (Per 21 CFR 801.109)

(Division Sign-Off)

Division of Reproductive, Abdominal,

and Radiological Devices

510(k) Number.

System:	Terason Model 2000 Porta 8IOL4	ble Uit	rasound	System					
	: Diagnostic ultrasound imagin	a or flu	id flow a	nalvsis of	the hun	nan body a	s follows:		
Clinical Applic			e of Ope		4.10 1101		10 10 HOWS.	•	
General	Specific	В	M	PWD	CWD	Calor	Comb.	Other	
(Track I Only)	(Tracks I & (II)				1	Dabb.	Modes	• • • •	
Ophthelmic	Ophthalmic	.I				1	7		
í	Felai ⁿ	I	Τ				 		
	Abdominal";						7		
	Intra-operative (Spec.)	N	N,	N		N	N.	N	
	Intra-operative (Neuro)								
Fetal	Laparoscopic Pediatric	ŧ	-			<u> </u>	 		
Imaging	Small Organ (Thyroid,	 	- 		 	├	 		
& Other	Breast, Testes, etc.)*:	£ .			ļ		1		
	Neonatal Cephalic:	1	 			 	 		
	Adult Cephalic":					 	 		
	Trans-rectal':		1			1	 		
	Trans-vaginal*:						1		
	Trans-urethral			_ [
	Trans-esoph. (non-Card.)						<u> </u>		
	Muscula-sket. (Convent.)				 				
	Musculo-skel. (Superfic)*: Intra-luminal	<u> </u>	 		ļ	 			
	Other (Specify)	-	╣	 		 	 		
	Cerdiac Adult	 	-	-		 	 		
Cardiac	Cardiac Fediatric		+			 	 		
	Trans-esoph (Cardiac)	 	+	+		 	╀───┤		
	Other (Specify)	L	╁			 	 		
Peripheral	Paripheral vessel*:		+	+			┼──		
/esset	Other (Specify)		+	 		├──-	╁╾╌╌╅		
Abdominal or Includes ultra: Includes ultra: Includes guid System uses System uses System uses System uses Includes Incl	sound guidance for placement gans and peripheral vessel. sound guidance for placement sound guidance of transvagina ance of amniocentesis, infertilit previously cleared under K912 previously cleared under K012 previously cleared under K010 previously cleared under K010 previously cleared under K010 no military field settings in addition WRITE BELOW THIS LINE	of nee I blops ty moni 505 wit 191. 1883. 191. ion to h	dies, cat y, infertiti toring of th 3 MHz nospital/c TINUE C	heters, cr ity monito fallicle de Model Li linic settir N ANOTH	ring of fi evelopme 3 (Linear ngs. HER PAC	ollicle deve ent. 7. SE IF NEE	elopment.	,	
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eratech Mode	1 810C4, 810L4, and 10LAP	4 Prob	cs			Page 4	3_A		
				Page 4.3-4					

System:	Terason Model 2000 Portable Ultrasound System

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows: Transducer: Mode of Operation Clinical Application Other PWD CWD Color Comb м В Specific General Modes* Dopp* (Tracks I & III) (Track I Only) Ophthalmic Ophthalmic Felal Abdominal N N, N, $\overline{\mathsf{N}}$ R N Intra-operative (Spec.) Intra-operative (Neuro) N N N N N N Laparoscopic Pediatric*: Fetal Small Organ (Thyrold, imaging & Other Breast, Testes, etc.) Neonatal Cephalic Adult Cephalic*: Trans-redial: Trans-vaginar Trans-urethral Trans-esoph. (non-Card.) Musculp-skel. (Convent.) Muscula-skel. (Superfic) Intra-lumina! Other (Specify) Cardiac Adult Cardiac Cardiac Pediatric Trans-esoph. (Cardiac) Other (Specify) Peripheral vessel*: Peripheral Other (Specify) Vessci

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System uses previously cleared under K012191.
System uses previously cleared under K010883.

* System uses previously cleared under K030191.

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(Division Sign-Off)

Division of Reproductive, Abdominal,

and Radiological Devices 510(k) Number .

12/09/04 TERATECH Corp. 510(k)

Teratech Model 8IOC4, 8IOL4, and 10LAP4 Probes

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